REMARKS

requested reconsideration in light of the foregoing amendments and following remarks is respectfully amendment can be found on page 8, line 9 of the application as filed. Reexamination and iso-alpha acids with the specific species of that genus without prejudice. Support for this essentially of" without prejudice. The claim has further been amended to replace the genus transition word in claim 13 has also been changed from "comprising" to "consisting throughout the specification and, for example, at page 1, line 20 or page 8, line 8. The with ostoeoarthritis or rheumatoid arthiritis.". Support for this amendment can be foun and the term "ostocoarthritis, rheumatoid arthiritis" has been replaced with "pain associate Claim 13 is under examination. With the this response, claim 13 has been amended

I. REJECTION UNDER 35 U.S.C. § 112, FIRST PARAGRAPH

Action, page 6, last line). Applicants respectfully disagree stating that "the claims do not find enablement from the instant specification" (Office that patients with osteoarthiritis had to have been tested. The Examiner concludes by (Office Action, page 4, lines 21-22). On page 6 of the Office Action, the Examiner states diseases/conditions were tested in vivo and no positive conclusions were ever drawn' acute pain (Office Action, page 3, lines 1-3). The Examiner states that "[t]he breath of the claims is enormous" (Office Action, page 4, line 13) and that "[n]one of the because the specification is not enabled for treating ostcoarthiritis, rheumatoid arthritis and with the enablement requirement. The Examiner states that "claim 13 is rejected. Claim 13 stands rejected under 35 U.S.C. § 112, first paragraph as failing to comply

analyzed eight factors in determining undue experimentation. Steel Corp. v. Sollac, 344 F.3d 1234, 1244 (Fed. Cir. 2003)). the specification, could practice the claimed invention without undue experimentation (ΔK The enablement requirement is satisfied when one skilled in the art, after reading MPEP 2164.01(a) The court in In re Wands

conclusion of nonenablement must be based on the evidence as a whole." examiner's analysis must consider all the evidence related to each of these factors, and any analysis of only one of the above factors while ignoring one or more of the others. The specifically states that it "is improper to conclude a disclosure is not enabling based on an

the breath of the claim is no longer as broad as the Examiner described it to be replacing the genus of iso alpha acids with its species. Accordingly, Applicants submit that with ostocoarthritis or rheumatoid arthiritis," replacing the transition word "comprising" and any of the Examiner's reasons for this rejection. As amended, the breath of the claim has been limited by replacing the term "ostocoarthritis, rheumatoid arthiritis" with "pain associate 13 solely to expedite the prosecution of the instant application and without acquiescing to Applicants disagree with the reasoning offered by the Examiner, they have amended claim namely, the breath of the claims and the existence of working examples. Although have disproportionately based his arguments on only two of the *In re Wands* factors. However, the Examiner, in support of the instant enablement rejection, appears to

its own be a determinative factor for whether or not the specification lacks enablement above, lack of working example is only one of the eight In re Wand's factors and cannot on certainly not a requirement of the Patent Office or the enablement standard. As mentioned showing in-vivo data may be the requirement of a regulatory agency such as FDA, it is of the enablement rejection and has mentioned at least four times on pages 4-6 of the Office Action. Applicants respectfully disagree with the Examiner and submit that although Lack of *in-vivo* studies is the next issue that the Examiner has focused on in support

Inc. v. Novo Nordisk NS, 108 F.3d 1361, 1367-68 (Fed. Cir. 1997) useful teaching, recognizing the stage of development of the technology." description in the specification must provide those skilled in the art with a specific and application of an unpredictable technology in the early stages of development, an enabling In deed, courts have recognized that "where, as here, the claimed invention is See Genentech,

that a skilled artisan is not enabled to practice the invention as claimed is unfounded of study that has been routinely done for different compounds and the Examiner's assertion of the specification, and a copy of which in included herewith). Indeed, WHMA is the type osteoarthritis or rheumatoid); and cited a reference on how to determine these activity Proc. Natl. Sci. USA 96:7563-68 (1999), incorporated by reference, on page 10, lines 10-13 levels or adjust the dosage range (e.g., through the William Harvey Human Modified NSAIDs that are commonly used in the treatment of acute pain or pain associated with teachings as to the dosage parameters; referred to appropriate activity levels (i.e., the IC50-Whole Blood Assay (WHMA) as described in detail in cited reference T.D. Warner et al., WHMA COX-2/COX-1 ratio) that is similar to that of other pain medications (e.g., The present application, rather than providing in-vivo data, has provided specific

undue experimentation. Applicants respectfully request the withdrawal of this rejection sufficient disclosure for one skilled in the art to practice the claimed invention without In re Wands factors, Applicants respectfully submit that the specification has provided Therefore, and in view of the reasons provided above and those of record for other

II. REJECTION UNDER 35 U.S.C. § 103(A)

range 5 mg to 1,000 mg claimed in claim 13. Office Action, page 8, third full paragraph bottle of beer produced in Todd contains an amount of isoalpha acids that falls within the evidenced by Medicinenet.com and About.com. In particular, the Examiner states that a al (US 3.354,219; hereinafter "Rigby") in view of Todd, Jr et al (US 5,041,300) as reasons Applicants respectfully traverse the rejection for the reasons of record and the following Claims 13 stands rejected under 35 U.S.C. § 103(a) as being anticipated by Rigby et

inhibitor having a COX-2/COX-1 ratio of about 0.23 to about 3.33 . . . , wherein the of "pharmaceutical composition consisting essentially of a therapeutic quantity of a COX-2 Applicants respectfully submit that claim 13 as amended is limited to administration

having a COX-2/COX-1 ratio of 0.23 to 3.33. COX-2 inhibitor, and beer is not known to be a pharmaceutical COX-2 inhibitor let alone beer composition because the claim is related to a composition 'consisting essentially' of a such, Applicants respectfully submit that claim 13 as amended does not read on the Todd's amount of the COX-2 inhibitor ranges from about 5 mg to about 1,000 mg per day." As

U.S.C. § 103(a) rejection of claim 13. does not render amended claim 13 obvious and respectfully request withdrawal of the 35 on the above reasons and the reasons of record, Applicants respectfully submit that Rigby of skill in the art to combine the references to produce the instant invention. in view of Todd, Jr et al (US 5,041,300) as evidenced by Medicinenet.com and About.com element of the amended claim nor provide any motivation or expectation of success for one rendered moot as the references cited, alone or in combination, neither teach each and every Therefore, Applicants respectfully submit that the ground for this rejection has been

III. DOUBLE PATENTING REJECTIONS

rejection. As such, Applicants respectfully request withdrawal of these rejections application and are therefore not a proper subject for a nonstatutory double patenting the above application and patent were both filed after the filing date of the instant patent No. 7279186, filed 01/09/2003. Applicants respectfully disagree on the basis that patenting over the claims of U.S. application No. 11409521, filed 4/21/2006, and U.S. Claim 13 has been rejected on the ground of ground of nonstatutory double

IV. CONCLUSION

respectfully requested submit that amended Claims 13 is in condition for allowance. Passage to issue is On the basis of the foregoing remarks and amendments, Applicants respectfully

Inventor: Kuhrts Application No. 10/008,778 Office Action Response

contact the undersigned at the telephone number provided below. If there are any questions regarding these remarks, the Examiner is encouraged to

tuture communications, to Deposit Account 50-1133. authorized to charge any fee under 37 C.F.R. § 1.17 applicable in this instant, as well as in Sunday) is included herewith. Pursuant to 37 C.F.R. § 1.136(a)(3), the Examiner is 2010 (the next successive business day after the due date of May 23, 2010, which fell on a A Request for a Three (3) Month Extension of Time, up to and including May 24,

appropriate length of time pursuant 37 C.F.R. § 1.136(a)(3) regardless of whether a separate petition is included. submission, as constructively incorporating a petition for extension of time for the requiring a petition for an extension of time under paragraph 1.136 for its timely Furthermore, such authorization should be treated in any concurrent or future reply

Respectively submitted,

MCDERMOTT WILL & EMERY LLP

Atabak R. Royacc, Ph.D. Agent for Applicants Registration No. 59,037

BST99 1650578-1.068911.0076

Date: May 24, 2010

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McDermott Will & Emery LLP

toxicity: A full in vitro analysis cyclo-oxygenase-2 are associated with human gastrointestinal Nonsteroid drug selectivities for cyclo-oxygenase-1 rather than

TIMOTHY D. WARNER**, FRANCESCO GIULIANO*, IVANA VOJNOVIC*, ANTOANETA BUKASA*, JANE A. MITCHELL‡ AND JOHN R. VANE*

The William Harvey Research Institute, St. Burtholomow, and the Royal Landon School of Netation; and Domstry, Charterhouse Squate, London, ECIM 68O United Kingdonn and Topperpount of Critical Care Medicine. The Royal Bompton (Strington and Superpount of Critical Care Medicine, The Royal Bompton (Strington and Coppensation SVM 50R), United Kingdonn.

Contributed by John R. Vanc. April 14, 1999

ABSTRACT
The beneficial actions of nonsteroid antiinflammatory drugs (NSALD) can be associated with inhibition
of cyclo-oxygenase (COX)-2 whereas their harmful side effects
are associated with inhibition of COX-1. Here we report data
from two related assay systems, the human whole blood assay
and a modified human whole blood assay (using human AS49
goells as a source of COX-2). This assay we refer to as the William
Harrey Modified Assay. Our aim was to make manningful
comparisons of both classical NSAIDs and newer COX-2,
selective compounds. These comparisons of the actions of >40
NSAIDs and discopropy! Huorophosphate, demonstrate a disribution of compound selectivities toward COX-1/2 selectivities in
with the risk of serious gastrointestinal complications. In conclusion, this full in vitro analysis of COX-1/2 selectivities in
human tissues clearly supports the theory that inhibition of
COX-1 underlies the gastrointestinal toxicity of NSAIDs in man.

therapeutic, anti-inflammatory effects of these agents are attrib-utable to their ability to unlihit (COX-2 (3). A number of subsequent analyses have been published demonstrating the patencies against COX-1 and COX-2 of a large number of NSAIDs and novel COX-2-selective inhibitors (see ref. 2). Alhas the advantage of using readily available human cells and taking into account the binding of NSA/IDs to human plasma proteins. However, thus far, there are no single studies published from isolated purified enzymes to intact cells, the assay most widely accepted is the human whole blood assay (4-7). This assay oxygenuse-1 (COX-1) is constitutive and present in, for example mechanism of action of the NSAIDs (1). As is now well appreciated, COX exists as two isoforms. In general terms, cycloderived from both the human whole blood assay (WBA) and use of NSAIDs in the patient population. Here we possible to determine the predictive nature of such assays for the appropriate assay system. Without such information, it is not family to inhibit COX-1 versus COX-2 on a common and that compare the relative abilities of all members of the NSAID though these analyses have used a wide range of assay systems, NSA IIDs correlate with their ability to inhibit COX-1 whereas the led some of us to the previous proposition that the side effects of in cells in vino and at inflammatory sites in vivo (see ref. 2). This (COX-2) is induced by proinflammatory cytokines and endolox in the endothelium, stomach and kidney whereas cyclo-oxygenase-2 (COX), and therefore prostaglandin production, is the common erative in laminatory joint diseases. Inhibition of cyclo-oxygenase choice in the treatment of rheumatic disorders and other degen-Nonsteroid anti-inflammatory drugs (NSAIDs) are among the most widely prescribed drugs worldwide, being the drugs of fitst present data

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human modified whole blood assay (WHMA) for >40 NSAIDs and COX-2-selective inhibitions. These data support the concept that inhibition of COX-1 is responsible for the serious gastrointestinal (G1) complications induced by NSAIDs in humans (8).

METHODS

Cell Culture, Human airway epithelial cells, A549 cells (European Collection of Animal Cell Cultures, ref. no. 86012804) were cultured in 96-well plates with DMEM supplemented with 10% fetal call serum and Legitamine (4 mM). To induce the expression of COX-2, A549 cells were exposed to interleukin-1β (10 ngm1-1) for 24 h (9).

Human Whole Blood Assay (WBA). Blood was collected by venupuncture into hepartin (19 units/ml) and then was aliquoted in 100-µd volumes into the individual wells of 96-well plates. For COX-1 assays, blood then was treated with test agents or vehicle casually 0.1% vol/vol dimethyl sulfoxdde) followed 60 min later by calcium ionephone, A23187 (50 µM). After 30 min, the plates were centrifuged (1,500 × g, 4°C, 5 min), and the plates were centrifuged (1,500 × g, 4°C, 5 min), and the plates were than the plates of the plates were span, and then 6 h later with hipopolysaccharide (10 µg/ml) bits test agents or vehicle. Incubation then was continued for a further 18 h, after with hipopolysaccharide (10 µg/ml) is unspeased in the plates were span, and the plasma was temoved and frozen. Concentrations of thromboxane (1'x) B₂ (as a measure of TXA₂ formulton and so COX activity) in samples from both protocols then were determined by radioimmunoassus. Data is reported as being from COX-1 and WBA-COX-2 protoceds.

William Harvey Human Modified Whole Blood Assay (WHMA). For assay of COX-1, experiments were conducted as above, and all COX-1 data were pooled. For assay of COX-2, the medium was removed from A549 cells, which had been exposed to interlenkin-1µ for the preceding 24 h, and human blood (100 µl) added together with test agents or vehicle. Sixy minutes later, A23187 (50 µM) was added, followed 30 min later by diedofenae (1 mM) to inhibit (>98%) the formation of prostanoids. The plates then were centrifuged, and plasma was removed (as above). Concentrations of prostaglandin E2 (PGE2) in samples then were determined by addisimmanosassay as a measure of the activity of COX-2 in the A549 cells. Data as reported as being from the WHMA-COX-2 protocols.

Materials. Radiolabeled [FHFRB, and [FHFOEs were obtained from Amersham, Celecowth, L-745.337, SC58125, and rofecowth were synthesized by Boehringer Ingelheim; 6-methoxy-2-naphlylacetic acid (oMNA) was a gift from SmithKline Beecham; discopropyl fluorophosphate was a gift from Merels-

Abhreviations: NSAID, nonsteroidal anti-inflammatery drug; COX, cyclo-oxygenase; WBA, whole blood assay; WHMA, William Harvey human modified whole blood assay; Tx, thromboxane; POE₂, prostagiandin E₃; 6MNA, 6-methoxy-2-napthylacetic acid; GI, gastrointestinal.

final. The whom reprint requests should be addressed, e-mail: t.d.warner@mds.qnnw.ac.uk.

Frosst Labs (Pointe Claire, PQ, Canada); tomoxiproie was a gitt from NicOx S.A. (Nies, France); ketorolae, meelofenamate, niflume acid, NS398, and valeryl salleylate were obtained from SPI Bio (Masy Codex, France); and sulfinder suffde was purchased from Affanti (Exeter, U.K.). All other compounds and reagents were obtained from Sigma.

Calculations. For each blood sample, the "control" formation of TsB₂ or PGE₂ was assessed as the mean of six determinations. For each experiment, the effects of the compounds were calculated and represented as percent of control by using the mean control value. Concentration response curves were fitted, and IC₂₀, and IC₂₀, values were derived, by using PRISM (GraphFad, San Diego). COX-1/WBA-COX-2 (WBA) and COX-1/WHMA-COX-2 (WHMA) selectivities were determined as the ratios of the IC₂₀ and IC₂₀, and IC₂₀

RESULTS

Prostanoid Production. In the presence of drug vehicle, the productions of prostanoids in the assay systems were COX-1, 32.3 ± 1.9 ngml⁻¹ TxB₂, WBA-COX-2, 12 ± 0.6 ngml⁻¹ TxB₃, and WHMA-COX-2, 41.8 ± 1.9 ngml⁻¹ PGE₂ (n = 24-31). In blood treated with aspirm and then incubated for 18 h in the absence of fipopolysaccharide, there was no detectable formation of TxB₂ or PGE₂.

Inhibitor Potencies, The agents tested readily divided into four groups in terms of their potencies as inhibitors of COX-1 and COX-2 (Table 1: Figs. 1–4). The first group consists of compounds that can produce full inhibition of both COX-1 and COX-2 with relatively poor selectivity. This group contained most of the currently used NSALDs, including, for instance, diciolenac, ibuprofen, naproxen, piroxicam, and sulindae (Fig. 1) as well as 6MNA. the active metabolite of nahumetone. Aspirin could not be assessed in the WBA-COX-2 assay because of its instability in whole blood but was active in the WHMA-COX-1 assay. Taken together with the COX-1 assay, our data demonstrated a selectivity of aspirin of ~4-fold loward COX-1. The second group contained compounds such as etodolae, meloxicam, and ninessulide, all of which show a preferential selectivity (Fig. 1). It must not be overlooked, however, that these compounds all have the potential to produce full inhibition of COX-1. Of interest, our data also indicate that exlections should be induded in this second group (Fig. 1). The third group contained compounds that inhibit COX-2, with only a very weak activity propyl fluorophosphate, L-745,37, NS598, and SCS8125 to selective agents (Fig. 2). The four third group contained compounds that inhibit cOX-2, which may contained as COX-1, and of the superintential compounds discopation of the subjective propyl fluorophosphate, L-745,37, NS598, and SCS8125 to such as many of the salicylates. As expected, this fourth group also included naburactone, which, untike its metabolite 6MNA, only produced weak inhibition of both COX is selected as many of the salicylates. As expected, this fourth group also included naburactone, which, untike its metabolitie 6MNA, only produced weak inhibition of both COX is selected.

DISCUSSION

Here, using simple assay systems, we have investigated the relative potencies as inhibitors of COX-1 and COX-2 of a wide range of NSAIDs as well as representatives of the newer COX-2 selective agents. In particular, however, we also included all of those agents for which good expidermological data of the risk of serious GI complications existed (8). This was a deliberate approach because, although some of these compounds were previously tested in other human whole blood assays (e.g., refs. 4–7), they have not been tested together within a single assay, system.

When comparing the potencies of NSAIDs against COX-1 and COX-2, ICov values are often used. However, there are assumptions underlying such an approach that are not necessarily correct in particular, as is clear from Figs. I and 2, the inhibitor curves are often not parallel. Thus, as the concentration of a NSAID varies, so does its relative potency. Second, NSAIDs are used thorspentically at doses that produce more than a 50% reduction

same time periods and in which the same stimulus is applied at the end of this incubation period, as for the matched COX-1 Insay system. Of interest, a number of the compounds tested appeared more potent in the WHMA-COX-2 hum the WBA-COX-2. This could be explained by variations in either the metabolism or the between the time courses of the incubations for testing inhibition of COX-1 and COX-2 (1 h w. 18 h) and hence, in the rate of prostanoid formation and so in the supply of arachidonic acid. The human whole blood plus A549 cell assay provides a system in which COX-2-containing cells are exposed to NSAIDs for the tolmetin (12), the steady-state plasma concentrations of these drugs, as well as the peak concentrations of aspirin (12), would produce average inhibitions in our assay systems of 82 ± 5% (COX-1), 74 ± 5% (WBA-COX-2), and 89 ± 2% (WHMA-COX-2). assay systems, or even to the binding characteristics of the NSAIDs to COX-2 (23). arachidonic acid within the cells expressing COX-2 in the two tively, it could be explained by different levels or sources of free the different time courses of the WBA and WHMA. Alternaplasma binding of compounds within the blood samples during human whole blood assay, there is a substantial difference acid both in vitro (21) and in vivo (22). Clearly, in the standard developed because the potencies of NSAIDs as inhibitors of more appropriate. In making these comparisons, we used data both from the WBA and from the WHMA. This second assay was against COX-1 and COX-2 at the ICs, value, therefore, appears naproxen (17), nimesulide (18), piroxicam (19), sulindac (20), and prostanoid formation are influenced by the supply of arachidonic COX-2) (n = 15). Comparison of the potencies of the NSAIDs ketorolac (13, 15), meclofenamate (12), meloxicam (16), (12, 13), fencyrofen (12), flurbiprofen (14), ketoprofen (12), established that, for dictofenac (10), stodolac (11), indomethacin in prostancid formation. Indeed, a survey of the literature When making our comparisons from the two assay systems we

of these compounds (Fig. 3) demonstrates that compounds associated with the greatest GI toxicity have the greatest COX-1 selectivity. These include tolments in udome thacin, ketoprofen (8), and, in particular, ketorolae. It is notable that we found ketorolae ketorolae is an extreme outlier both in our assay system and in NSA1Ds (25). Clearly, this is in keeping with the idea that COX-1 inhibition underlies the serious GI complications of NSAIDs. because this compound is to be the most COX-1 selective of all of the NSAIDs we tested produce serious GI complications by significantly inhibiting the activity of COX. Further comparison of the COX-1 selectivities not included azapropazone in any of our subsequent analyses). Group I (see Table I) contained all of the NSAIDs included in this analysis. This is consistent with the idea that NSAIDs ketoprofen, and 11-tolmetin, with azapropazone last. (We have 1-ibuprofen, 2-dictofenac, 3-diffunisal, 4-fenoprofen, 5-ass 6-sulindac, 7-naproxen, 8-indomethacin, 9-piroxicam, zone) were ordered for their association with serious compilea-toms. The order of the NSA IDs. from least to most damaging, was 1-ibuprofen. 2-dictofenac, 3-diffunisal, 4-fenoprofen, 5-aspirju. tween 1985 and 1994 (8) in which 11 NSAIDs (plus azapropamost complete recent studies is a meta-analysis of reports behas, therefore, been examined in a number of studies. One of the groupings of NSAIDs to epidemiological studies of NSAID-induced GI toxicity. This is an area of particular interest, for NSAIDs cause serious gastric damage leading to hospitalization in some 100,000 patients per year in the U.S. alone (24). The relationship between NSAID use and serious GI complications compounds that appeared to be only weak inhibitors of COX-1 and COX-2 (Table 1; Fig. 3). It is of interest to compare these groups; (f) compounds capable of producing full inhibition of both COX-1 and COX-2 with poor selectivity; (f) compounds capable of producing full inhibition of COX-1 and COX-2 with preference toward COX-2; (if) compounds that strongly inhib-ted COX-2 with only weak activity against COX-1; and (p)) found that the agents tested could be divided into four main ~5× more gastrotoxic than other

and WHMA-COX-2 Potencies of all compounds tested as inhibitors of prostanoid formation determined in the COX-1 assay. WBA-COX-2,

	10%	- 1"	1Cm. 1Ch.	1C80.	ICso.	Co. ICa.	WBA	WHMA	VBW	IC 36 Fattos	WBA ICso	A WIINA
Compound	M	M	MA	Z	Z	L.N.	COX-1	COX-1	COX-1	COX-1	COX-I	COX-1
6MNA	42	130	146	580	n.d.	n.d.	3.5	n.d.	4.5	n.d.	27	n.d.
Aspirin	1.7	8.0	V 106	001 v	7.5	30	> 100	4.	>100	3.8	34	23
Carproten	0.087	. 50	4	75	n.d.	n,d,	50	n.d.	3.9	n.d.	2.5	n.d.
Diclotenac	0.075	0.1	0.038	0.27	0.020	0.23	0.5	0.3	0.27	0.23	0	9
Fenoproten	1 (1) 14.	23	-	00	5.9	24	1.2	1.7	1.3	0.3	26	81
Fishersten	3076	- 8	n V	2 6	n.d.	n.d.	13.1	n. d.	1.0	n d	Į,	n.d.
Duprofen	7.6	ź.	40	674	3 p.//	160	3	10	14	Ch.	يرا د سر	27
Indomerhacin	0.013	0.45	5 i	5.0	0.13	120	800	5 1.0	- 1	4 F.	ž T	2 t
Ketoprofen	0.047	1.0	2.9	2.2	0.2.4	6.0	2 3	A .	22:	5 (ا در پاس	0 1
Ketorolac	6100016	0.0034	0.926	4.0	0.075	5	453	395	176	204	اند خلفه	×
Meclofenamate	0.22	9.0	0.7	x.C	0.2	- -	3.2	0.91	2.7	0 -	22	- 6
Metenamic acid	13	100	2.9	>100	1.3	√ 120	0.11	0.049	. !		. :	. :
Naproxen	9.3	003	28	260	S.	330	3.0	ω 30	14	3.0	20	22
Niffumic acid	ß	77	5.4	i.	,,,t	74	0.22	0.43	0.45	0	1 7	16
Piroxicam	2.4	15	7.9	31	0.17	7.0	ندا ندأ	0.1	(3	0.47	1.7	<u>.</u>
Sulindac sulphide	1.9	38	55	100	1.21	2.	29	0.64	1,0	0.29	20	5
Suprofen	C	3.0	8.7	56	۵ نیا	100	7.7	7.3	19	33	30	26
Tenidap	0.081	5.0	2.9	57	n.d.	n.d.	35.2	n.d.	2.6	n.d.	21	n.d.
Tolmetin	0.35	5.0	0.82	43	1.3	13	2.3	32 50	8.0	2.6	28	21
l'omoxiprol	7.6	35	20	×	0.32	1.3	2.7	0.042	24	0.37	63	12
Zomepirac	0.43	2.0	0.81	6.0	0.096	2.0	1 9	0.22	3.0	C.3	23	17
Celexocib	1.2	28	0.83	6.0	0.34	3.0	0.7	0.3	0.21	0.11	vc	7
Etodolac	5	69	2.2	8.0	0.94	30	0.2	0 1	0.12	0.043	0	Uni
Meloxicam	5.7	22	2.1	7	0.23	2.0	0.37	0.040	0.32	160'0	-13	6
Nimesulide	10	4	1.9	7.0	0.39	7.0	61.0	0.038	0.17	0.17	7	x
Diisopropyi					;			:				
745 337		7 1	0 0 0	2 :	- 9.	1	1000	1000	10.00	10.01		ī
Z9198	o 6	200	0 00	- 1	20.5		0.00	20.01		A9.91	- 1	·
Rofecoxib		9 60	0.81	^ ·	0.03	n :	2000	0.0001		1000	٠ ١	v +
SC58125	> 100 ×	>100	2.0	ŏ	n.d.	n.d.	>0.01	n.d.	<0.01	n.d.	1	n,o
5-Aminosalicylic												
acid	¥	>1000	16	>1000	n.d.	n.d.	0.15	n.d.	,	n.c.		n.d.
Ampyrone		270	203	0001	85	070	3.7	1.5	3.7	2.5	4	19
Diflunisal		530	50	140	134	400	0.1	1.2	0.26	0.75	ç	4
Nahumetone		> 1000	> 1000	>1000	290	>1000		•	•	,	ı	ŧ
Paracetamol		V 100	÷ co	003	0.4	>100		,	ì		,	•
Resveratroi		>100	9	001V	n.d.	n.d.	1.3	n.d		,	,	n.d.
Salicin		V 100	V100	>100	n.d.	n.d.	,	n.d.	٠		1	n.d.
Sancylaidenyde		>100	>100	V (00	200	n.d.	,	n.d.		,		n.d.
Sodium salicylare		00000	34440	000101	482	450W	6.9	0.10	13	0.92	5	Ĭĸ
Sulfasalazine		6400	2507	8300	5 4	n.d.	8.0	n,d.	1.3	n.d	3.50	n.d.
Sulindac		>100	> 100	>100	58	-100	1			4	,	,
Tamoxiten		V 100	Se	√ 100	n.d.	n,d.	6.4	n.G.	,		,	n,d
Tick-padine	52 V	¥ 68	47	>100	n.d.	n.d.	0.9	# G			4	n.d.
		CKIL	2.3	¥ 69	n.d.	n.d.	0.053	n.d.		n.d.		n C
Valeryl salicylate	24											

produce full inhibition of both COX-1 and COX-2 with poor COX-2 selectivity, (second) compounds that can produce full inhibition of COX-1 and COX-2. With poor COX-2 selectivity, (second) compounds that appear to be only weak inhibitors of COX-1 and COX-2. Shown are potencies (independent Fig. and ICM, voltees) of compounds against COX-1. WBA-COX-2 and WHMA-COX-2. Selectivities of compounds towards (COX-1 were determined as ICM, and ICM, voltees) of compounds against COX-1, wBA-COX-2, and WHMA-COX-2. Selectivities of compounds towards COX-1 were determined as ICM, and ICM products for both WBA-COX-2/COX-1 and WHMA-COX-2/COX-1. Ranking of compounds as inhibitors of COX-2 relative to COX-1 are based on ordering of ICM ratios, higher ranking numbers are associated with increased selectivity towards COX-1, n.d., not done.

Because all of the compounds contained within group 1 have the potential to produce full inhibition of both COX-1 and COX-2, their associated risk of producing GI toxicity can be strongly influenced by dose. This can be cadily appreciated by reference to Fig. 4. Here we have displayed the extent of COX-1 inhibition produced by individual NSA IDs at concentrations that cause 80% inhibition of COX-2. This analysis essentially provides the answer to the important question. If a NSA ID is used at levels sufficient to inhibit COX-2 by 80%, i.e., to produce some therapeutic effect, by how much will COX-1 be inhibited? As can be seen, the classical NSA IDs produce inhibitions of ~80% or more.

This implies that, even for a drug such as dictofenac, which is >4-told selective for COX-2 in terms of C_{low} values, therapeuticully relevant selectivity will be very difficult to achieve; i.e., the concentration of dictofenac necessary to produce 80% inhibition of COX-1. To extend this line of reasoning, it is also clear that, when relative selectivities differ by only slight amounts, other variables, such as ingested dose and plasma half-life, will have a particular influence on NSA1D toxicity (20). This may well be especially true for proxicam, which we did not find in our assays to be notably COX-1-selective despite its well established G1 toxicity. Phoxi-

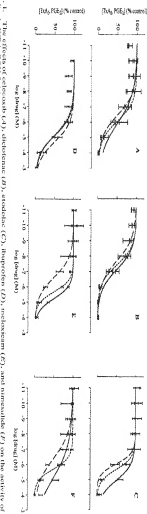
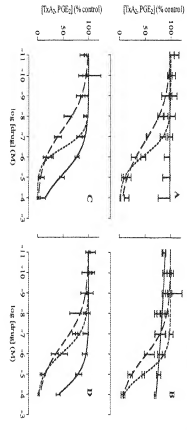


Fig. 1. The effects of celecoxib (4), diclotense (θ), etodolac (C), ibuptofen (D), meloxicam (E), and nimesulide (F) on the activity of COX-1 (solid line). WBA-COX-2 (short dashed line), and WHMA-COX-2 (long dashed line). Results are expressed as percent of control and are represented as mean \pm SEM: (n = 5-8).

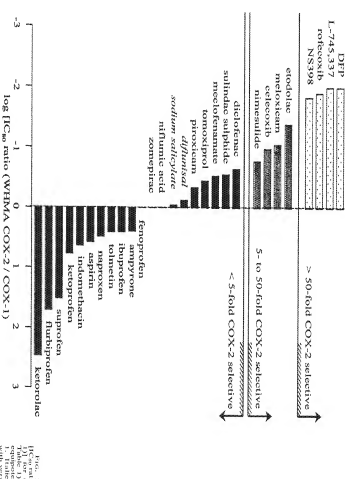
cara, however, has a much longer elimination half-life (30 to 70 h) (19) than other NSA IDs, and plasma half-life has been previously correlated with GI toxicity (27).

compounds in this group are capable of inhibiting this isoform of that increasing the dosage of these agents could readily increase Gl toxicity due to inhibition of COX-1 because all of the an improved GI toxicity profile. It must be remembered, however, mcloxicam (28, 29)] show that these preferential compounds have to inhibit COX-2 by 80% produce only 25% inhibition of COX-1. example, the concentrations of etodolac and meloxicam sufficient tivity of these compounds could be usefully exploited. COX (Fig. 1). Despite the sparse epidemiological data, controlled trials [e.g., for COX-1. Possibly more importantly, Fig. 4 implies that the selecpounds with between 5- and 50-fold selectivity for COX-2 over COX-2 inhibitors. In Fig. Tie second grouping of NSAIDs consists of preferential we have classified these as com-For

It is interesting that, in our assays, eelecoxib was found to be a member of the preferential group of COX-2 inhibitors. This is in contrast to data derived by using recombinant human COX-1 and COX-2 from broken insect cells. In this system, celecoxib is between 155- and 3,200-fold selective for COX-2 over COX-1 (23). This difference may be autributable to the fact that celecoxib inhibition of both COX-1 and COX-2 is initially competitive with respect to substrate and is characterized by similar affinity for COX-1 and COX-2. There is a second, slow, time-dependent briding of celecoxib to COX-2 but not COX-1 that may well produce the selectivity seen in other assay systems (23). It is currently not eleter why celecoxib does not demonstrate such selectivity in either the WBA or WHMA. It is unlikely that these assay systems in some way delay the time-dependent binding of celecoxib to COX-2. For instance, in the isolated human enzyme assays, this secondary binding takes place in secondar stather than minutes (23), and the WHMA assay included a preincubsation period of 60 mm, and the WBA included a 24-b incubation period.



Pio. 2. The effects of disopropyl fluorophosphate (A). 1-745,337 (B), NS398 (C), and collecasts (D) on the activity of COX-2 (elso il disting), WBA-COX-2 (elso il disting), wBBAwHMA-COX-2 (led unquisted wHMA-COX-2 (led unquisted with a converse of control and agre represented as mean ± SEM (n = 5-8).



Our data also reinforce the concept that compounds within group 3 that inhibit COX-2 with only a very weak activity against COX-1 will produce few serious GI complications when used in the general population. As is clear from both the direct inhibitor curves (Fig. 2) and the derived data (Figs. 3 and 4), these compounds produce very little effect on COX-1 and should have a large therapeutic window. There are preliminary reports that rofecoxib has a low GI toxicity, but, until appropriate comparative clinical trads have been completed, no firm conclusions can be drawn (30). Furthermore, it must be remembered that studies in animals (31) suggest that when used in the presence of existing GI damage. COX-2-selective inhibitors rulght slow the repair process in man due to reductions in the production of protective COX-2 products (32).

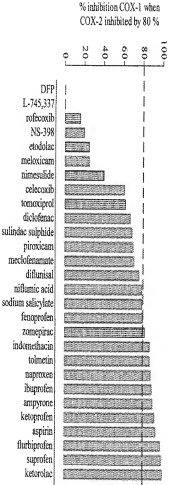
Group 4 contains weak inhibitors of COX-1 and COX-2 for which reliable data with regard to inhibition of COX-1 and

Fig. 3. Determinable log IF.G., 3. Determinable log IF.G. patie (WBA_CON_2/CON_2).

If.G. patie (WBA_CON_2/CON_2).

If for all agents assayed (see Table 1). The "0 line" indicates equipotency, i.e., an IC.g. ratio of 1. Italies indicate compounds with very low potency.

COX-2 could not be derived. These compounds are not, therefore, displayed in Figs. 3 and 4. Clearly, however, the weak ability of the group 4 compounds to inhibit prostanoid production explains their general lack of, or very low, GI toxicity Sodium saleylate, for example, only caused inhibition of prostanoid formation at concentrations for in excess of those achieved in tho (13) and in accordance with its relatively low GI toxicity (33). As expected, this fourth group also contained nabunetone whereas its series metabolite, 6MAA (34), was a member of the first group. This classification is in accordance with the results of Patrigian et al. (4) who found that oral dosing of indumentone at I guertaly for 7 days reduced COX-1 activity in the WBA by 70%. The plasma concentration of drug achieved with such dosing (34) would correlate with the activity of 6MNA but not nabunetone, which we report here. As a cautionary remark to other mystigators, we would like to note that we also tested as additional gators, we would like to note that we also tested as additional



n 80% Analysis of the percent inhibition of COX-I seen when COX-2 (WHMA) is inhibited by 80%. The dotted line indicates equiactivity % inhibition of COX-I.

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COX-1 (WBA). active found habumelone to be essentially inactive and 6MNA to be the activity of and selectivity of nabumetone and 6MNA. variations in supply may explain some of the confusion regarding assay systems similar to that of nabumetone. Possibly such were found to be essentially inactive, with potencies in the various samples of "6MNA" supplied from commercial sources. These all In conclusion, we have conducted a full and careful with a selectivity at the ICso values of 4.5-fold toward our ra

analysis of COX-1/2 selectivities for a large range of NSA IDs and COX-2-selective compounds. The distribution of potenties of these agents as inhibitors of COX-1 relative to COX-2 supports. gastrointestinal toxicity of NSAIDs. our earlier premise (3) that inhibition of COX-1 underlies the

supported by a grant from Bochringer Ingelheim. J.A.M. is T.D.W. holds a British Heart Foundation Lectureship (BS/95003), and A.M. is a Wellcome Career Development fellow. This work was

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